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(57) Abstract

Catalase enzymes derived from bacterial for the genera Alcaligenes (Delaya) and MicroscUla are disclosed. The enzymes are produced from native or recombinant host cells and can be utilized to destroy or detect hydrogen peroxide, e.g., in production of glyoxylic acid and in glucose sensors, and in processes where hydrogen peroxide is used as a bleaching or antibacterial agent, e.g., in contact lens cleaning, in bleaching steps in pulp and paper preparation and in the pasteurization of dairy products.

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CATALASES

Field of the Invention

This invention relates generally to enzymes and more specifically to catalases and polynucleotides encoded such catalases, including methods of use.

5 Background

This invention relates to newly identified polynucleotides, polypeptides encoded by such polynucleotides, the use of such polynucleotides and polypeptides, as well as the production and isolation of such polynucleotides and polypeptides.

More particularly, the polynucleotides and polypeptides of the present invention have been putatively identified as catalases.

Generally, in processes where hydrogen peroxide is a by-product, catalases can be used to destroy or detect hydrogen peroxide, e.g., in production of glyoxylic acid and in glucose sensors. Also, in processes where hydrogen peroxide is used as a bleaching or antibacterial agent, catalases can be used to destroy residual hydrogen peroxide, e.g. in contact lens cleaning, in bleaching steps in pulp and paper preparation and in the pasteurization of dairy products. Further, such catalases can be used as catalysts for oxidation reactions, e.g., epoxidation and hydroxylation.

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Summary of the Invention

In accordance with one aspect of the present invention, there are provided novel enzymes, as well as active fragments, analogs and derivatives thereof.

In accordance with another aspect of the present invention, there are 5 provided isolated nucleic acid molecules encoding the enzymes of the present invention including mRNAs, cDNAs, genomic DNAs as well as active analogs and fragments of such enzymes.

In accordance with yet a further aspect of the present invention, there is provided a process for producing such polypeptides by recombinant techniques 10 comprising culturing recombinant prokaryotic and/or eukaryotic host cells, containing a nucleic acid sequence of the present invention, under conditions promoting expression of said enzymes and subsequent recovery of said enzymes.

In accordance with yet a further aspect of the present invention, there are also provided nucleic acid probes comprising nucleic acid molecules of sufficient 15 length to specifically hybridize to a nucleic acid sequence of the present invention.

In accordance with yet a further aspect of the present invention, there is provided a process for utilizing such enzymes, or polynucleotides encoding such enzymes, for in vitro purposes related to scientific research, for example, to generate probes for identifying similar sequences which might encode similar enzymes from 20 other organisms by using certain regions, i.e., conserved sequence regions, of the nucleotide sequence.

In accordance with yet a further aspect of the present invention, there is provided antibodies to such catalases. These antibodies are as probes to screen libraries from these or other organisms for members of the libraries which could have 25 the same catalase activity or a cross reactive activity.

In another embodiment, the invention provides a method for catalyzing an oxidation reaction comprising contacting a substrate with an effective amount of an enyzme selected from the group consisting of an amino acid sequence set forth in SEQ ID NOS: 7 or 9, thereby catalyzing an oxidation reaction. Another method of 30 the invention includes the detection and/or destruction of hydrogen peroxide in a

sample comprising contacting the sample with an effective amount of an enzyme having an amino acid sequence set forth in SEQ ID NO:7 or SEQ ID NO:9, and detecting the presence of hydrogen peroxide in the sample. Hydrogen peroxide acts as a substrate for catalases, thus, either the detection and/or the destruction of hydrogen peroxide is achieved by combining a sufficient amount of the catalases of the invention with a sample or material suspected of containing hydrogen peroxide.

These and other aspects of the present invention should be apparent to those skilled in the art from the teachings herein.

Brief Description of the Drawings

The following drawings are illustrative of an embodiment of the invention and are not meant to limit the scope of the invention as encompassed by the claims.

Figure 1 shows the full-length DNA sequence and the corresponding deduced amino acid sequence for *Alcaligenes (Deleya) aquamarinus* Catalase - 64CA2.

Figure 2 shows the full-length DNA sequence and the corresponding deduced amino acid sequence for *Microscilla furvescens* Catalase 53CA 1.

Detailed Description of Preferred Embodiments

In order to facilitate understanding of the following description and examples which follow certain frequently occurring methods and/or terms will be described.

The term "isolated" means altered "by the hand of man" from its natural state; i.e., if it occurs in nature, it has been changed or removed from its original environment, or both. For example, a naturally occurring polynucleotide or a polypeptide naturally present in a living animal in its natural state is not "isolated",

but the same polynucleotide or polypeptide separated from the coexisting materials of its natural state is "isolated", as the term is employed herein. For example, with respect to polynucleotides, the term isolated means that it is separated from the nucleic acid and cell in which it naturally occurs.

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As part of or following isolation, such polynucleotides can be joined to other polynucleotides, such as DNAs, for mutagenesis, to form fusion proteins, and for propagation or expression in a host, for instance. The isolated polynucleotides, alone or joined to other polynucleotides such as vectors, can be introduced into host cells, in culture or in whole organisms. Introduced into host cells in culture or in whole organisms, such polynucleotides still would be isolated, as the term is used herein, because they would not be in their naturally occurring form or environment. Similarly, the polynucleotides and polypeptides may occur in a composition, such as a media formulation (solutions for introduction of polynucleotides or polypeptides, for example, into cells or compositions or solutions for chemical or enzymatic reactions which are not naturally occurring compositions) and, therein remain isolated polynucleotides or polypeptides within the meaning of that term as it is employed herein.

The term "ligation" refers to the process of forming phosphodiester bonds

between two or more polynucleotides, which most often are double stranded DNAs.

Techniques for ligation are well known to the art and protocols for ligation are described in standard laboratory manuals and references, such as, for instance,

Sambrook et al., MOLECULAR CLONING, A LABORATORY MANUAL, 2nd Ed.;

Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1989).

The term "gene" means the segment of DNA involved in 4producing a polypeptide chain; it includes regions preceding and following the coding region (leader and trailer) as well as intervening sequences (introns) between individual coding segments (exons).

A coding sequence is "operably linked to" another coding sequence when RNA polymerase will transcribe the two coding sequences into a single mRNA, which is then translated into a single polypeptide having amino acids derived from both coding sequences. The coding sequences need not be contiguous to one another so long as the expressed sequences ultimately process to produce the desired protein.

"Recombinant" enzymes refer to enzymes produced by recombinant DNA techniques; i.e., produced from cells transformed by an exogenous DNA construct

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encoding the desired enzyme. nSynthetic" enzymes are those prepared by chemical synthesis.

A DNA "coding sequence of" or a "nucleotide sequence encoding" a particular enzyme, is a DNA sequence which is transcribed and translated into an enzyme when placed under the control of appropriate regulatory sequences.

"Plasmids" are designated by a lower case "p" preceded and/or followed by capital letters and/or numbers. The starting plasmids herein are either commercially available, publicly available on an unrestricted basis, or can be constructed from available plasmids in accord with published procedures. In addition, equivalent plasmids to those described are known in the art and will be apparent to the ordinarily skilled artisan.

"Digestion" of DNA refers to catalytic cleavage of the DNA with a restriction enzyme that acts only at certain sequences in the DNA. The various restriction enzymes

used herein are commercially available and their reaction conditions, cofactors and other requirements were used as would be known to the ordinarily skilled artisan. For analytical purposes, typically 1 μg of plasmid or DNA fragment is used with about 2 units of enzyme in about 20 μl of buffer solution. For the purpose of isolating DNA fragments for plasmid construction, typically 5 to 50 μg of DNA are digested with 20 to 250 units of enzyme in a larger volume. Appropriate buffers and substrate amounts for particular restriction enzymes are specified by the manufacturer. Incubation times of about 1 hour at 37.C are ordinarily used, but may vary in accordance with the supplier's instructions. After digestion the reaction is electrophoresed directly on a polyacrylamide gel to isolate the desired fragment.

Size separation of the cleaved fragments is performed using 8 percent polyacrylamide gel described by Goeddel et al., *Nucleic Acids Res.*, 8:4057 (1980).

"Oligonucleotides" refers to either a single stranded polydeoxynucleotide or two complementary polydeoxynucleotide strands which may be chemically synthesized. Such synthetic oligonucleotides have no 5' phosphate and thus will not ligate to another oligonucleotide without adding a phosphate with an ATP in the

presence of a kinase. A synthetic oligonucleotide will ligate to a fragment that has not been dephosphorylated.

"Ligation" refers to the process of forming phosphodiester bonds between two double stranded nucleic acid fragments (Maniatis, T., et al., Id., p. 146). Unless otherwise provided, ligation may be accomplished using known buffers and conditions with 10 units of T4 DNA ligase ("ligase") per 0.5 μg of approximately equimolar amounts of the DNA fragments to be ligated.

Unless otherwise stated, transformation was performed as described in Sambrook and Maniatis, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, 1989.

In accordance with an aspect of the present invention, there are provided isolated nucleic acids (polynucleotides) which encode for the mature enzyme having the deduced amino acid sequence of Figure 1 (SEQ ID NO: 7).

In accordance with another aspect of the present invention, there are provided isolated nucleic acids (polynucleotides) which encode for the mature enzyme having the deduced amino acid sequence of Figure 2 (SEQ ID NO: 9).

In accordance with another aspect of the present invention, there is provided an isolated polynucleotide encoding the enzyme of the present invention. The deposited material is a genomic clone comprising DNA encoding an enzyme of the present invention. As deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852, USA, the deposited material is assigned ATCC Deposit No.

The deposit has been made under the terms of the Budapest Treaty on the International Recognition of the Deposit of Micro-organisms for Purposes of Patent

25 Procedure. The clone will be irrevocably (without restriction or condition) released to the public upon the issuance of a patent. This deposit is provided merely as convenience to those of skill in the art and is not an admission that a deposit would be required under 35 U.S.C. §112. The sequence of the polynucleotide contained in the deposited material, as well as the amino acid sequence of the polypeptide encoded

30 thereby, are controlling in the event of any conflict with any description of sequences

herein. A license may be required to make, use or sall the deposited material, and no such license is hereby granted.

The polynucleotides of this invention were originally recovered from a genomic gene library derived from two sources. The first, *Alcaligenes (Delaya)*5 aquamarinus, is a β-Proteobacteria. It is a gram-negative rod that grows optimally at 26° C and pH 7.2. The second, *Microscilla furvescens*, is a Cytophagales (Bacteria) isolated from Samoa. It is a gram-negative rod with gliding motility that grows optimally at 30° C and pH 7.0.

With respect to Alcaligenes (Delaya) aquamarinus, the protein with the closest amino acid sequence identity of which the inventors are currently aware is the Microscilla furvescens catalase (59.5 % protein identity; 60 % DNA identity). The next closest is a Mycobacterium tuberculosis catalase (KatG), with a 54 % protein identity.

With respect to *Microscilla furvescens*, the protein with the closest amino acid sequence identity of which the inventors are currently aware is catalase I of *Bacillus stearothermophilas*, which has a 69% amino acid identity.

Accordingly, the polyoucleotides and enzymes encoded thereby are identified by the organism from which they were isolated. Such are sometimes referred to below as "64CA2" (Figure 1 and SEQ ID NOS: 6 and 7) and "53CA1" (Figure 2 and SEQ ID NOS: 8 and 9).

One means for isolating the nucleic acid molecules encoding the enzymes of the present invention is to probe a gene library with a natural or artificially designed probe using art recognized procedures (see, for example: Current Protocols in Molecular Biology, Ausubel F.M. et al. (EDS.) Green Publishing Company Assoc.

25 and John Wiley Interscience, New York, 1989, 1992). It is appreciated by one skilled in the art that the polynucleotides of SEQ ID NOS: 6 and 8, or fragments thereof (comprising at least 12 contiguous nucleotides), are particularly useful probes. Other particularly useful probes for this purpose are hybridizable fragments of the sequences of SEQ ID NOS: 6 and 8 (i.e., comprising at least 12 contiguous nucleotides).

With respect to nucleic acid sequences which hybridize to specific nucleic acid sequences disclosed herein, hybridization may be carried out under conditions of reduced stringency, medium stringency or even stringent conditions. As an example of oligonucleotide hybridization, a polymer membrane containing immobilized denatured nucleic acids is first prehybridized for 30 minutes at 45°C in a solution consisting of 0.9 M NaCl, 5.0 mM NaH₂PO₄, pH 7.0, 5.0 mM Na₂EDTA, 0.5% SDS, 10X Denhardt's, and 0.5 mg/mL polyriboadenylic acid. Approximately 2 X 10⁷ cpm (specific activity 4-9 X 10⁸ cpm/ug) of ³²p end-labeled oligonucleotide probe are then added to the solution. After 1216 hours of incubation, the membrane is washed for 30 minutes at room temperature in 1X SET (150 mM NaCl, 20 mM Tris hydrochloride, pH 7.8, 1 mM Na₂EDTA) containing 0.5% SDS, followed by a 30 minute wash in fresh 1X SET at (Tm less 10°C) for the oligonucleotide probe. The membrane is then exposed to auto-radiographic film for detection of hybridization signals.

Stringent conditions means hybridization will occur only if there is at least 90% identity, preferably at least 95 % identity and most preferably at least 97% identity between the sequences. Further, it is understood that a section of a 100 bps sequence that is 95 bps in length has 95% identity with the 1090 bps sequence from which it is obtained. See J. Sambrook et al., Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory (1989) which is hereby incorporated by reference in its entirety. Also, it is understood that a fragment of a 100 bps sequence that is 95 bps in length has 95% identity with the 100 bps sequence from which it is obtained.

As used herein, a first DNA (RNA) sequence is at least 70% and preferably at least 80% identical to another DNA (RNA) sequence if there is at least 70% and preferably at least a 80% or 90% identity, respectively, between the bases of the first sequence and the bases of the another sequence, when properly aligned with each other, for example when aligned by BLASTN.

The present invention relates to polynucleotides which differ from the reference polynucleotide such that the differences are silent, for example, the amino acid sequence encoded by the polynucleotides is the same. The present invention also

relates to nucleotide changes which result in amino acid substitutions, additions, deletions, fusions and truncations in the polypeptide encoded by the reference polynucleotide. In a preferred aspect of the invention these polypeptides retain the same biological action as the polypeptide encoded by the reference polynucleotide.

The polynucleotides of this invention were recovered from genomic gene libraries from the organisms identified above. Gene libraries were generated from a Lambda ZAP II cloning vector (Stratagene Cloning Systems). Mass excisions were performed on these libraries to generate libraries in the pBluescript phagemid. Libraries were generated and excisions were performed according to the protocols/methods hereinafter described.

The polynucleotides of the present invention may be in the form of RNA or DNA, which DNA includes cDNA, genomic DNA, and synthetic DNA. The DNA may be double stranded or single-stranded, and if single stranded may be the coding strand or non-coding (anti-sense) strand. The coding sequences which encodes the mature enzymes may be identical to the coding sequences shown in Figures 1-2 (SEQ ID NOS: 6 & 8) or may be a different coding sequence which coding sequence, as a result of the redundancy or degeneracy of the genetic code, encodes the same mature enzymes as the DNA of Figures 12 (SEQ ID NOS: 6 & 8).

The polynucleotide which encodes for the mature enzyme of Figures 1-2

(SEQ ID NOS: 7 & 9) may include, but is not limited to: only the coding sequence for the mature enzyme; the coding sequence for the mature enzyme and additional coding sequence such as a leader sequence or a proprotein sequence; the coding sequence for the mature enzyme (and optionally additional coding sequence) and non-coding sequence, such as introns or noncoding sequence 5' and/or 3' of the coding sequence for the mature enzyme.

Thus, the term "polynucleotide encoding an enzyme (protein)" encompasses a polynucleotide which includes only coding sequence for the enzyme as well as a polynucleotide which includes additional coding and/or non-coding sequence.

The present invention further relates to variants of the hereinabove described polynucleotides which encode for fragments, analogs and derivatives of the enzymes having the deduced amino acid sequences of Figures 1-2 (SEQ ID NOS: 7 & 9). The variant of the polynucleotide may be a naturally occurring allelic variant of the polynucleotide or a nonnaturally occurring variant of the polyoucleotide.

Thus, the present invention includes polynucleotides encoding the same mature enzymes as shown in Figures 1-2 (SEQ ID NOS: 7 & 9) as well as variants of such polynucleotides which variants encode for a fragment, derivative or analog of the enzymes of Figures 1-2 (SEQ ID NOS: 7 & 9). Such nucleotide variants include deletion variants, substitution variants and addition or insertion variants.

As hereinabove indicated, the polynucleotides may have a coding sequence which is a naturally occurring allelic variant of the coding sequences shown in Figures 1-2 (SEQ ID NOS: 6 & 8). As known in the art, an allelic variant is an alternate form of a polynucleotide sequence which may have a substitution, deletion or addition of one or more nucleotides, which does not substantially alter the function of the encoded enzyme. Also, using directed and other evolution strategies, one may make very minor changes in DNA sequence which can result in major changes in function.

Pragments of the full length gene of the present invention may be used as hybridization probes for a cDNA or a genomic library to isolate the full length DNA and to isolate other DNAs which have a high sequence similarity to the gene or similar biological activity. Probes of this type preferably have at least 10, preferably at least 15, and even more preferably at least 30 bases and may contain, for example, at least 50 or more bases. In fact, probes of this type having at least up to 150 bases or greater may be preferably utilized. The probe may also be used to identify a DNA clone corresponding to a full length transcript and a genomic clone or clones that contain the complete gene including regulatory and promotor regions, exons and introns. An example of a screen comprises isolating the coding region of the gene by using the known DNA sequence to synthesize an oligonucleotide probe. Labeled oligonucleotides having a sequence complementary or identical to that of the gene or

portion of the gene sequences of the present invention are used to screen a library of genomic DNA to determine which members of the library the probe hybridizes to.

It is also appreciated that such probes can be and are preferably labeled with an analytically detectable reagent to facilitate identification of the probe. Useful reagents include but are not limited to radioactivity, fluorescent dyes or enzymes capable of catalyzing the formation of a detectable product. The probes are thus useful to isolate complementary copies of DNA from other sources or to screen such sources for related sequences.

The present invention further relates to polynucleotides which hybridize to the hereinabove-described sequences if there is at least 70%, preferably at least 90%, and more preferably at least 95% identity between the sequences. (As indicated above, 70% identity would include within such definition a 70 bps fragment taken from a 100 bp polynucleotide, for example.) The present invention particularly relates to polynucleotides which hybridize under stringent conditions to the hereinabove-described polynucleotides. As herein used, the term "stringent conditions" means hybridization will occur only if there is at least 95 % and preferably at least 97% identity between the sequences. The polyoucleotides which hybridize to the hereinabove described polynucleotides in a preferred embodiment encode enzymes which either retain substantially the same biological function or activity as the mature enzyme encoded by the DNA of Figures 1-2 (SEQ ID NOS: 6 & 8). In referring to identity in the case of hybridization, as known in the art, such identity refers to the complementarily of two polynucleotide segments.

Alternatively, the polynucleotide may have at least 15 bases, preferably at least 30 bases, and more preferably at least 50 bases which hybridize to any part of a polynucleotide of the present invention and which has an identity thereto, as hereinabove described, and which may or may not retain activity. For example, such polynucleotides may be employed as probes for the polynucleotides of SEQ ID NOS: 6 & 8, for example, for recovery of the polyoucleotide or as a diagnostic probe or as a PCR primer.

Thus, the present invention is directed to polynucleotides having at least a 70% identity, preferably at least 90% identity and more preferably at least a 95% identity to a polynucleotide which encodes the enzymes of SEQ ID NOS: 7 & 9 as well as fragments thereof, which fragments have at least 15 bases, preferably at least 30 bases, more preferably at least 50 bases and most preferably fragments having up to at least 150 bases or greater, which fragments are at least 90% identical, preferably at least 95% identical and most preferably at least 97% identical to any portion of a polynucleotide of the present invention.

The present invention further relates to enzymes which have the deduced amino acid sequences of Figures 1-9 (SEQ ID NOS: 28-36) as well as fragments, analogs and derivatives of such enzyme.

The terms "fragment,n nderivative" and "analog" when referring to the enzymes of Figures 1-9 (SEQ ID NOS. 28-36) means enzymes which retain essentially the same biological function or activity as such enzymes. Thus, an analog includes a proprotein which can be activated by cleavage of the proprotein portion to produce an active mature enzyme.

The enzymes of the present invention may be a recombinant enzyme, a natural enzyme or a synthetic enzyme, preferably a recombinant enzyme.

The fragment, derivative or analog of the enzymes of Figures 1-2 (SEQ ID NOS: 7 & 9) may be (i) one in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code, or (ii) one in which one or more of the amino acid residues includes a substituent group, or (iii) one in which the mature enzyme is fused with another compound, such as a compound to increase the half-life of the enzyme (for example, polyethylene glycol), or (iv) one in which the additional amino acids are fused to the mature enzyme, such as a leader or secretory sequence or a sequence which is employed for purification of the mature enzyme or a proprotein sequence. Such fragments, derivatives and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.

The enzymes and polynucleotides of the present invention are preferably provided in an isolated form, and preferably are purified to homogeneity.

The present invention also relates to vectors which include polynucleotides of the present invention, host cells which are genetically engineered 5 with vectors of the invention and the production of enzymes of the invention by recombinant techniques.

Host cells are genetically engineered (transduced or transformed or transfected) with the vectors of this invention which may be, for example, a cloning vector such as an expression vector. The vector may be, for example, in the form of a 10 plasmid, a phage, etc. The engineered host cells can be cultured in conventional nutrient media modified as appropriate for activating promoters, selecting transformants or amplifying the genes of the present invention. The culture conditions, such as temperature, pH and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

The polynucleotides of the present invention may be employed for producing enzymes by recombinant techniques. Thus, for example, the polynucleotide may be included in any one of a variety of expression vectors for expressing an enzyme. Such vectors include chromosomal, nonchromosomal and synthetic DNA sequences, e.g., derivatives of SV40; bacterial plasmids; phage DNA; baculovirus; 20 yeast plasmids; vectors derived from combinations of plasmids and phage DNA, viral DNA such as vaccinia, adenovirus, fowl pox virus, and pseudorabies. However, any other vector may be used as long as it is replicable and viable in the host.

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The appropriate DNA sequence may be inserted into the vector by a variety of procedures. In general, the DNA sequence is inserted into an appropriate 25 restriction endonuclease site(s) by procedures known in the art. Such procedures and others are deemed to be within the scope of those skilled in the art.

The DNA sequence in the expression vector is operatively linked to an appropriate expression control sequence(s) (promoter) to direct mRNA synthesis. As representative examples of such promoters, there may be mentioned: LTR or SV40 30 promoter, the E. coli. lac or trp, the phage lambda P_L promoter and other promoters

known to control expression of genes in prokaryotic or eukaryotic cells or their viruses. The expression vector also contains a ribosome binding site for translation initiation and a transcription terminator. The vector may also include appropriate sequences for amplifying expression.

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In addition, the expression vectors preferably contain one or more selectable marker genes to provide a phenotypic trait for selection of transformed host cells such as dihydrofolate reductase or neomycin resistance for eukaryotic cell culture, or such as tetracycline or ampicillin resistance in *E. coli*.

The vector containing the appropriate DNA sequence as hereinabove

described, as well as an appropriate promoter or control sequence, may be employed to transform an appropriate host to permit the host to express the protein.

As representative examples of appropriate hosts, there may be mentioned: bacterial cells, such as *E. coli*, *Streptomyces, Bacillus subtilis*; fungal cells, such as yeast; insect cells such as *Drosophila S2* and *Spodoptera Sf9*; animal cells such as 15 CHO, COS or Bowes melanoma; adenoviruses; plant cells, *etc.* The selection of an appropriate host is deemed to be within the scope of those skilled in the art from the teachings herein.

More particularly, the present invention also includes recombinant constructs comprising one or more of the sequences as broadly described above. The constructs comprise a vector, such as a plasmid or viral vector, into which a sequence of the invention has been inserted, in a forward or reverse orientation. In a preferred aspect of this embodiment, the construct further comprises regulatory sequences, including, for example, a promoter, operably linked to the sequence. Large numbers of suitable vectors and promoters are known to those of skill in the art, and are commercially available. The following vectors are provided by way of example; Bacterial: pQE70, pQE60, pQE-9 (Qiagen), pBluescript II KS(Stratagene), ptrc99a, pKK223-3, pDR540, pRIT2T (Pharmacia); Eukaryotic: pXT1, pSG5 (Stratagene) pSVK3, pBPV, pMSG, pSVL SV40 (Pharmacia). However, any other plasmid or vector may be used as long as they are replicable and viable in the host.

Promoter regions can be selected from any desired gene using CAT

(chloramphenicol transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, apt, lambda PR, PL and trp. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art.

In a further embodiment, the present invention relates to host cells containing the above-described constructs. The host cell can be a higher eukaryotic cell, such as a mammalian cell, or a lower eukaryotic cell, such as a yeast cell, or the host cell can be a prokaryotic cell, such as a bacterial cell. Introduction of the construct into the host cell can be effected by calcium phosphate transfection, DEAE-Dextran mediated transfection, or electroporation (Davis, L., Dibner, M., Battey, I., Basic Methods in Molecular Biology, (1986).

The constructs in host cells can be used in a conventional manner to

15 produce the gene product encoded by the recombinant sequence. Alternatively, the
enzymes of the invention can be synthetically produced by conventional peptide
synthesizers.

Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described by Sambrook et al., Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor, N.Y., (1989), the disclosure of which is hereby incorporated by reference.

25 Transcription of the DNA encoding the enzymes of the present invention by higher eukaryotes is increased by inserting an enhancer sequence into the vector. Enhancers are cisacting elements of DNA, usually about from 10 to 300 bp that act on a promoter to increase its transcription. Examples include the SV40 enhancer on the late side of the replication origin bp 100 to 270, a cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and

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adenovirus enhancers.

Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, e.g., the ampicillin resistance gene of E. coli and S. cerevisiae TRP1 gene, and a promoter derived from a highlyexpressed gene to direct transcription of a downstream

5 structural sequence. Such promoters can be derived from operons encoding glycolytic enzymes such as 3-phosphoglycerate kinase (PGK), a-factor, acid phosphatase, or heat shock proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences, and preferably, a leader sequence capable of directing secretion of translated enzyme.

10 Optionally, the heterologous sequence can encode a fusion enzyme including an N-terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product.

Useful expression vectors for bacterial use are constructed by inserting a structural DNA sequence encoding a desired protein together with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector and to, if desirable, provide amplification within the host. Suitable prokaryotic hosts for transformation include *E. coli, Bacillus subtilis, Salmonella typhimurium* and various species within the genera Pseudomonas, Streptomyces, and Staphylococcus, although others may also be employed as a matter of choice.

As a representative but nonlimiting example, useful expression vectors for bacterial use can comprise a selectable marker and bacterial origin of replication derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and pGEM1 (Promega Biotec, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed.

Following transformation of a suitable host strain and growth of the host

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strain to an appropriate cell density, the selected promoter is induced by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period.

Cells are typically harvested by centrifugation, disrupted by physical or 5 chemical means, and the resulting crude extract retained for further purification.

Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents, such methods are well known to those skilled in the art.

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Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described by Gluzman, Cell, 23: 175 (1981), and other cell lines capable of expressing a compatible vector, for example, the C127, 3T3, CHO, HeLa and BHK cell lines. Mammalian expression vectors will comprise 15 an origin of replication, a suitable promoter and enhancer, and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements.

The enzyme can be recovered and purified from recombinant cell cultures by methods including ammonium sulfate or ethanol precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, afflinty chromatography, hydroxylapatite chromatography and lectin chromatography. Protein refolding steps can be used, as 25 necessary, in completing confi-uration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps.

The enzymes of the present invention may be a naturally purified product. or a product of chemical synthetic procedures, or produced by recombinant 30 techniques from a prokaryotic or eukaryotic host (for example, by bacterial, yeast,

higher plant, insect and mammalian cells in culture). Depending upon the host employed in a recombinant production procedure, the enzymes of the present invention may be glycosylated or may be non-glycosylated. Enzymes of the invention may or may not also include an initial methionine amino acid residue.

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Antibodies generated against the enzymes corresponding to a sequence of the present invention can be obtained by direct injection of the enzymes into an animal or by administering the enzymes to an animal, preferably a nonhuman. The antibody so obtained will then bind the enzymes itself. In this manner, even a sequence encoding only a fragment of the enzymes can be used to generate antibodies 10 binding the whole native enzymes. Such antibodies can then be used to isolate the enzyme from cells expressing that enzyme.

The term "antibody," as used herein, refers to intact immunoglobulin molecules, as well as fragments of immunoglobulin molecules, such as Fab, Fab', (Fab')₂, Fv, and SCA fragments, that are capable of binding to an epitope of an 15 endoglucanase polypeptide. These antibody fragments, which retain some ability to selectively bind to the antigen (e.g., an endoglucanase antigen) of the antibody from which they are derived, can be made using well known methods in the art (see, e.g., Harlow and Lane, *supra*), and are described further, as follows.

- (1) A Fab fragment consists of a monovalent antigen-binding fragment of an 20 antibody molecule, and can be produced by digestion of a whole antibody molecule with the enzyme papain, to yield a fragment consisting of an intact light chain and a portion of a heavy chain.
- (2) A Fab' fragment of an antibody molecule can be obtained by treating a whole antibody molecule with pepsin, followed by reduction, to yield a molecule consisting 25 of an intact light chain and a portion of a heavy chain. Two Fab' fragments are obtained per antibody molecule treated in this manner.
 - (3) A (Fab')₂ fragment of an antibody can be obtained by treating a whole antibody molecule with the enzyme pepsin, without subsequent reduction. A (Fab'), fragment is a dimer of two Fab' fragments, held together by two disulfide bonds.

- (4) An Fv fragment is defined as a genetically engineered fragment containing the variable region of a light chain and the variable region of a heavy chain expressed as two chains.
- (5) A single chain antibody ("SCA") is a genetically engineered single chain molecule
 containing the variable region of a light chain and the variable region of a heavy chain, linked by a suitable, flexible polypeptide linker.

As used in this invention, the term "epitope" refers to an antigenic determinant on an antigen, such as an endoglucanase polypeptide, to which the paratope of an antibody, such as an endoglucanase-specific antibody, binds.

10 Antigenic determinants usually consist of chemically active surface groupings of molecules, such as amino acids or sugar side chains, and can have specific threedimensional structural characteristics, as well as specific charge characteristics.

For preparation of monoclonal antibodies, any technique which provides antibodies produced by continuous cell line cultures can be used. Examples include the hybridoma technique (Kohler and Milstein, *Nature*, 256:495-497, 1975), the trioma technique, the human B-cell hybridoma technique (Kozbor et al., *Immunology Today* 4:72, 1983), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole et al., in *Monoclonal Antibodies and Cancer Therapy*, Alan R. Liss, Inc., pp. 77-96, 1985).

Techniques described for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce single chain antibodies to immunogenic enzyme products of this invention. Also, transgenic mice may be used to express humanized antibodies to immunogenic enzyme products of this invention.

Antibodies generated against an enzyme of the present invention may be used in screening for similar enzymes from other organisms and samples. Such screening techniques are known in the art, for example, one such screening assay is described in Sambrook and Maniatis, Molecular Cloning: A Laboratory Manual (2d Ed.), vol. 2:Section 8.49, Cold Spring Harbor Laboratory, 1989, which is hereby incorporated by reference in its entirety.

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The present invention will be further described with reference to the following examples; however, it is to be understood that the present invention is not limited to such examples. All parts or amounts, unless otherwise specified, are by weight.

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Example 1

Production of the Expression Gene Bank

An *E. coli* catalase negative host strain CAT500 was infected with a phage solution containing sheared pieces of DNA from *Alcaligenes (Deleya) aquamarinus* in pBluescript plasmid and plated on agar containing LB with ampicillin (100 ~g/mL), methicillin (80 ~g/mL) and kanamycin (100 ~g/mL) according to the method of Hay and Short (Hay, B. and Short, J., *J. Strategies*, 5:16, 1992). The resulting colonies were picked with sterile toothpicks and used to singly inoculate each of the wells of 96-well microtiter plates. The wells contained 250 ,uL of SOB media with 100 ~g/mL ampicillin, 80 ~g/mL methicillin, and (SOB Amp/Meth/Kan). The cells were grown overnight at 37°C without shaking. This constituted generation of the "SourceGeneBankn; each well of the Source GeneBank thus contained a stock culture of *E. coli* cells, each of which contained a pBluescript plasmid with a unique DNA insert. Same protocol was adapted for screening catalase from *Microscilla furvescens*.

Example 2

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Screening for Catalase Activity

The plates of the Source GeneBank were used to multiply inoculate a single plate (the "Condensed Plate") containing in each well 200 µL of SOB Amp/Meth/Kan. This step was performed using the High Density Replicating Tool (HDRT) of the Beckman Biomek with a 1 % bleach, water, isopropanol, air-dry sterilization cycle in between each inoculation. Each well of the Condensed Plate thus contained 4 different

pBluescript clones from each of the source library plates. Nine such condensed plates were prepared and grown for 16h at 37°C.

One hundred (100) µL of the overnight culture was transferred to the white polyfiltronic assay plates containing 100 µL Henes/well. A 0.03% solution of hydrogen peroxide was made in 5 % Triton and 20 µL of this solution was added to each well. The plates were incubated at room temperature for one hour. After an hour, 50 ,µL of 120 mM 3-(p-hydroxyphenyl)-propionic acid and 1 unit of horseradish peroxidase were added to each well and the plates were incubated at room temperature for 1 hour. To quench the reaction, 50 ,µL of 1 M Tris-base was added to each well. The wells were excited on a fluorometer at 320 nm and read at 404 nm. A low value signified a positive catalase hit.

Example 3 Isolation and Purification of the Active Clone

In order to isolate the individual clone which carried the activity, the

Source GeneBank plates were thawed and the individual wells used to singly inoculate a new plate containing SOB Amp/Meth/Kan. As above the plate was incubated at 37°C to grow the cells, and assayed for activity as described above. Once the active well from the source plate was identified, the cells from the source plate were streaked on agar with LB/Amp/Meth/Kan and grown overnight at 37°C to obtain single colonies. Eight single colonies were picked with a sterile toothpick and used to singly inoculate the wells of a 96well microtiter plate. The wells contained 250 pL of SOB Amp/Meth/Kan. The cells were grown overnight at 37°C without shaking. A 100 µL aliquot was removed from each well and assayed as indicated above. The most active clone was identified and the remaining 150 µL of culture was used to streak an agar plate with LB/Amp/Meth/Kan. Eight single colonies were picked, grown and assayed as above. The most active clone was used to inoculate 3mL cultures of LB/Amp/Meth/Kan, which were grown overnight. The plasmid DNA was isolated from the cultures and utilized for sequencing.

Example 4

Expression of Catalases

DNA encoding the enzymes of the present invention, SEQ ID NOS: 7 and 9, were initially amplified from a pBluescript vector containing the DNA by the PCR technique using the primers noted herein. The amplified sequences were then inserted into the respective pQE vector listed beneath the primer sequences, and the enzyme was expressed according to the protocols set forth herein. The 5' and 3' oligonucleotide primer sequences used for subcloning and vectors for the respective genes are as follows:

- 10 Alcaligenes (Deleya) aquamarinus catalse: (pQET vector)
 - 5' Primer

CCGAGAATTCATTAAAGAGGAGAAATTAACTATGAATAACGCATCCGCTG AC EcoRI (SEQ ID NO:1)

3 ' Primer CGGAAAGCTTTTACGACGCGACGTCGAAACG HindI I I (SEQ ID

15 NO:2)

Microscilla furvescens catalase: (pQET vector)

5' Primer

CCGAGAATTCATTAAAGAGGAGAAATTAACTATGGAAAATCACAAACACT CA EcoRI (SEQ ID NO:3)

20 3' Primer CGAAGGTACCTTATTTCAGATCAAACCGGTC Kpnl (SEQ ID NO:4)

The restriction enzyme sites indicated correspond to the restriction enzyme sites on the bacterial expression vector indicated for the respective gene (Qiagen, Inc. Chatsworth, CA). The pQET vector encodes antibiotic resistance (Ampr), a bacterial origin of replication (ori), an IPTG-regulatable promoter operator (P/O), a ribosome

25 binding site (RBS), a 6-His tag and restriction enzyme sites.

The pQET vector was digested with the restriction enzymes indicated. The amplified sequences were ligated into the respective pQET vector and inserted in

frame with the sequence encoding for the RBS. The native stop codon was incorporated so the genes were not fused to the His tag of the vector. The ligation mixture was then used to transform the E. cold strain UM255tpREP4 (Qiagen, Inc.) by electroporation. UM255/pREP4 contains multiple copies of the plasmid pREP4, 5 which expresses the lacl repressor and also confers kanamycin resistance (Kanr). Transformants were identified by their ability to grow on LB plates and ampicillin/kanamycin resistant colonies were selected. Plasmid DNA was isolated and confirmed by restriction analysis. Clones containing the desired constructs were grown overnight (O/N) in liquid culture in LB media supplemented with both Amp 10 (100 u μ /ml) and Kan (25 u μ /ml). The O/N culture was used to inoculate a large culture at a ratio of 1:100 to 1:250. The cells were grown to an optical density 600 (O.D. 600) of between 0.4 and 0.6. IPTG ("Isopropyl-B-D-thiogalacto pyranosiden") was then added to a final concentration of 1 mM. IPTG induces by inactivating the lacl repressor, clearing the P/O leading to increased gene expression. Cells were grown an extra 3 to 4 hours. Cells were then harvested by centrifugation. The primer sequences set out above may also be employed to isolate the target gene from the

deposited material by hybridization techniques described above.

- 25 -

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What Is Claimed Is:

- Substantially pure catalase having an amino acid sequence of SEQ ID NO:7 or SEQ ID NO:9
- 2. An isolated polynucleotide sequence encoding a catalase of claim 1.
- 3. An isolated polynucleotide selected from the group consisting of:
 - a) SEQ ID:6 or SEQ ID NO:8;
 - b) SEQ ID:6 or SEQ ID NO:8, wherein T can also be U;
 - c) nucleic acid sequences complementary to a) and b); and
 - d) fragments of a), b), or c) that are at least 15 bases in length and that will selectively hybridize to DNA which encodes the amino acid sequences of SEQ ID Nos:7 or 9, respectively.
- 4. The polynucleotide of claim 2, wherein the polynucleotide is isolated from a prokaryote.
- 5. An expression vector including the polynucleotide of claim 2.
- 6. The vector of claim 5, wherein the vector is a plasmid.
- 7. The vector of claim 5, wherein the vector is a virus-derived.
- 8. A host cell transformed with the vector of claim 5.
- 9. The host cell of claim 8, wherein the cell is prokaryotic.
- 10. Antibodies that bind to the polypeptide of claim 1.

- 11. The antibodies of claim 10, wherein the antibodies are polyclonal.
- 12. The antibodies of claim 10, wherein the antibodies are monoclonal.
- 13. An enzyme comprising a member selected from the group consisting of:
 - a) an enzyme comprising an amino acid sequence which is at least 70% identical to the amino acid sequence set forth in SEQ ID NO:7 or SEQ ID NO:9; and
 - b) an enzyme which comprises at least 30 amino acid residues to an enzyme of a).
- 14. A method for producing an enzyme comprising growing a host cell of claim 8 under conditions which allow the expression of the nucleic acid and isolating the enzyme encoded by the nucleic acid.
- 15. A process for producing a cell comprising: transforming or transfecting the cell with the vector of Claim 5 such that the cell expresses the polypeptide encoded by the DNA contained in the vector.
- 16. A method for catalyzing an oxidation reaction comprising contacting a substrate with an effective amount of an enyzme selected from the group consisting of an amino acid sequence set forth in SEQ ID NOS: 7 or 9, thereby catalyzing an oxidation reaction.
- 17. A method for detection or destruction of hydrogen peroxide in a sample comprising contacting the sample with an effective amount of an enzyme having an amino acid sequence set forth in SEQ ID NO:7 or SEQ ID NO:9, and detecting the presence of hydrogen peroxide in the sample.

FIGURE 1

Alcaligenes (Deleya) aquamarinus Catalase - 64CA2

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461	Asp	Leu	Ile	Trp	Gln	Aep	Pro	Ile	Pro	Al=	GIA	Aen	Thr	yab	Tyr	суa	gju	Glu	Val	VAI	460
1441	AAG	CAG	AAA	ATT	GCA	CAA	AGT	GGC	CIG	AGC	ATT	AGT	GAG	DTA	CTC	TCC	ACC	CCI	TGG	CAC	1500
481	Larm	aln	Tara	Ile	Ala	Gla	Ser	Glv	Leu	Ser	Ile	Ser	Glu	Mat	Val	Ser	Thr	Ala	Trp	Asp	500
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1561	TTC	CCC	CCA	CAG	AAC	CAO	TGG	CAG	GGC	AAC	GAG	cca	CLAG	CGC	CIG	GCG	-		-	AGC	
521	Leu	Ala	Pro	Gln	Asn	Glu	Trp	Gln	Gly	Asn	Glu	Pro	Glu	Arg	Leu	X1.a	Lye	Val	Leu	sar	540
1621	GTC	TAC	GAG	CAG	ATC	TCT	GCC	GAC	ACC	GGC	GCT	AGC	ATC	aca	CIAC	GIG	ATC	GII	CLG	GCC	1680
541	Val	Tyr	<u>Olu</u>	Gla	11e	Ser	Ala	Asp	Thr	Gly	Ala	Ser	Ile	Ala	Asp	Val	Ile	Val	Leu	Ala	560
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1681	GGT	300	CTT B	ccc	ATC	GAG	222	acc	GCO	AAA	GCA	GCA	GGT	TAC	CAT	crc	CGC	GTT	CCC	TTC	1740
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581	Lau	Lys	CIA	ErA	Gly	Asp	λla	Thr	Ala	Glu	Met	Thr	хвр	ALE	Авр	241	Pne	W14	*10	Leu	800
1801	GAG																				1860
601	Glu	Pro	Leu	Ala	Asp	Gly	Phe	Arg	Asn	Trp	Gln	Lys	Lys	Glu	Tyr	Val	Val	Lys	Pro	Glu	620
1361	CAG	ATG	cre	CTG	GAT	CST	ေ	CAG	CTG	ATG	GGC	TTA	ACC	GGC	CCS	GAA	ATG	ACC	GTG	CLC	1920
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661	λøp	Сув	Glu	Gly	Gln	Leu	Thr	Aan	Хар	Pha	Phe	Val	λsn	Leu	Thr	yab	Met	ath	Asn	ser	680
2041	TGG	AAG	cca	GTA	GGT	AGC	AAC	GCC	TAC	CYY	RTC	CGC	CAC	CGC	AAG	ACC	COT	GCC	CLC	AAG	2100
681	Tro	Lva	Pro	Val	Gly	Ser	Asn	Ala	Tyr	Glu	Ile	Arg	Asp	Arg	Lys	Thr	Gly	Ale	Val	Lys	700
		-1-		,	•				-												
2101	TIECT	ACC	ccc	TCG	CCC	GTG	GAT	CTG	GTA	TIT	GGT	TCC	AAC	TCG	CTA	CTG	CGC	TCT	TAC	GCA	2160
	T	The	A1 -	Ser	1	Val	Ago	Lau	Val	Phe	alv	Ser	Asn	Ser	Leu	Leu	λrg	Ser	Tyr	Ala	720
701	rrb	inr	VII	382	~£4		~•h				1								•		
				GCC	<b></b> -		C++	886	occ	020	210	<b>TT</b> C	arc.	ACIB	CAC	TTC	GTC.	acc	GCC	TGG	2220
2161	UAA _^	ard	IAC	GCC Ala	-AG	-	mv.	***	0	~~	~~·	Db-	Vel	1	200	ph-	Vel	11-	A1 =	Tre	740
721	Glu	Val	TYT	Ala	GIU	Asp	veb	ASI.	ork	OLU	-ye	21/4		~. 9	~~b			<b>-</b>	·		
														<b></b> -							
2221				ATG												162					
741	Thr	Lye	Val	Het	Aan	Als	Yeb	λrg	Phe	Yeb	Val	Ala	Ser	End	75	• •					

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#### FIGURE 2

#### Microscilla furvescens Catalago 53CA1

	~ 1	• •	~ ~			~~	CAC					, vc	. 1X	. ~~		^ ^^	- AL	1 40	c cu	~ ^^	A TUC	• • •
1	Мо	t G	u As	in H	le L	.ys	His	Set	Gly	. Sei	: Se	r Th	r Ty	r As	n Th	r Aa	n Th	r Gl	y al	y Ly	в сув	20
61	cc	T	TAC	c oc	IA (I	ют	TCO	cn		cu	, AG1	raci	A GG	т оо	c <b>g</b> g	C AC	CAA	A AA	C AG	n da	T TGO	120
21	Pr	0 21	e T:	- 01	уG	ly	Ser	Leu	Lye	Qln	841	: Al	G1;	y G1	A OT	y Th	r Ly	s Aas	n Ar	g As	p Trp	40
121	TC	G CC	C AA	.C A1	'a c	TC	AAC	CTC	GGC	ATC	TTA	CGC	CM	, CA	r rci	. TC		. TCC	CA.	. cc	A AAC	180
41																					o Asn	60
		_																	_			
181																					AAG Lys	240 80
	•						-,-				• • • • • • • • • • • • • • • • • • • •	-/-	-,-								-,-	
241																					TAT	300
81	Ase	Le	u Al	# WI	a L	au i	Met	Thr	Aap	Ser	GIN	Asp	Trp	Trp	Pro	N1=	Авр	lyr	OLY	Hla	Tyr	100
301	GGC	: cc	TT	e TT	T A7	CA (	ccc	DTA	GCG	TGG	CAC	AGC	GCC	ago	ACC	TAC	CGT	ATC	GGT	CAT	GGC	360
101	aly	Pr	Pho	Ph:	e Il	e 3	Arg	Met	Ala	Trp	His	Ser	Ala	Gly	Thr	Tyr	Arg	11e	Gly	Aop	Gly	120
361	CGT	. co.	r age	: aa	rod	ic 1	rcc	aac	TCA	CAG	cac	TTC	aca	ccr	CTC	AAT	λGC	TGG	CCA	GAC	AAT	420
121																Adn						140
141																CAA Gln						480 160
					. –,			•							-•		•	•	•	_	•	
481																CIG						540
161	Ile	Ser	III	Al:	. As	pΙ	.eu	Met	Il.	Leu	Thr	GIY	Aen	Val	XII	Leu	Old	Thr	Met	στλ	Phe	180
\$41	XXX	ACT	111	GG1	TI	TG	ica i	cat	GGC	AGA	GCA	CAT	σīλ	TGG	CAG	CCI	caa	CYA	GAT	GTA	TAC	600
181	Lye	The	Phe	aly	2h	• A	le	Gly	αlγ	Arg	<b>7</b> ]=	Asp	Val	Trp	Glu	Pro	Glu	Gļu	Aap	Val	Tyr	200
€01	TGG	GGA	GCA	GAR	, AC	cc	aa :	TGG	CTG	GGA	GAC	AAG	œc	TAT	GAA	CCT	GAC	CSA	ರ್ಷ	CIC	GAA	660
201																Gly						220
661				-		~ ~	~~	~ .	3 TV-1	ccz	<del></del>	h ~~~	~~ ~	OT.	B 5 C	CCC	<b>GB B</b>	CG N	ccc	110	ccc	720
221																Pro						240
721																III Phe						780 260
241	~y s	PLO	wab	FEU	. 11	9 ^	14 /			nu y	veb	-11	~	414			<b>4.</b>				,,,,,	400
																TTC						640
261	Asn	λap	Glu	Glu	Th	r V	al J	lla:	L⊕U.	Ile .	λla	GΙΆ	GIĀ	H1#	Thr	Phe	GIÅ	Lys	Thr	Hla	GIA	280
841	CCT	GCC	GAT	aca	GA	G A	AA 1	TAT	org (	SGC	CGA	CAG	CCI	GCC	GCC	GCA	GGT	ATT	GAA	CAA	ATG	900
281	Ala	Ala	yab	Al=	Gl	u L	ys 7	Δr ,	V=1 (	sly .	Arg	Glu	Pro	Alu	Ala	Ala	Gly	Ila	Glu	Glu	Met	300
901	AGC	cro	caa	TGG	, AA	A A.	AC /	NCC '	TAC (	aac .	ACC	gga.	CAC	GGT	aca	GAT	ACC	ATC	ACC	AGT	GGA	960
301																Asp						320
961			~~~		-					~~~	. —	<b></b> 1	<b>*</b>	100			-	-	<b>~</b>			1020
321																						1020 340
			•																			
021																						1080
341	ens.	ork	TAX	OLU	ιη	y U	tu I	∍ <b>e</b> u ⊺	AIT I	у•	-=I	PEO.	~14	GIY	~7=	• Y *	- III	Þ	~ y 5	210	υγ•	360
081																						1140
361	Aap	Glv	Ala	Glv	Ale		1v 2	thr :	Ile :	Pro /	Asp.	Ala	Hie	λap	Pro	Ser	Ly=	Sat	Him	Ala	Pro	380

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1141		N T C	<b>ст</b> с	a (~T	a CG	GAC	cra	aca	CIG	cac	DTA	GAC	CCT	GAT	TAC	GAA	AAA	ATT	τcτ	CGA	1200
1141	Pho i	MAF	Lau	Thr	Thr	Aan	Leu	Ala	Leu	Arg	Het	Asp	Pro	Aep	τγτ	alu	Lye	Ila	Ser	Arg	400
1201	caa	TAC	TAT	CLAA	AAC	ca	CAT	CAG	III	GCA	CAT	GCT	TTC	oca	AAA	Q CX	ĹŒ	TAC	AAA	CTO	1360
401	Arg	Tyr	Туг	Glu	Asn	Pro	Asp	alu	Ph=	Alm	λ≠р	<b>X1</b> =	Spe	W/F	Lys	Ala	Trp	Tyr	Lye	Leu	420
1261	ACA	CXC	ACA	GAT	ATG	GGA	CCA	AAG	CTC	COC	TAC	CTG	QQA	CCA	GYY.	QIQ.	CCI	CXG	an	CAC	1320
421	Thr	Ki•	Arg	Asp	Met	σlγ	Pro	Lys	Val	Arg	Ty	Leu	gly	Pro	Olu	Val	Pro	@Tu	OLu	Asp	440
												~~ ~		~~~	OTA	CAC	a a a	BAC.	GD T	ATT	1360
1321	CIC	ATC	TGG	CAA	GAC	CCI	ATA	CCA	CAT	GT.	AGC	CAI	CCI	tan	V-1	and and	O) ii	Aan	Ann	710	460
441	Leu	Ile	Trp	Gln	qaA	Pro	Ile	Pro	Aap	ATI	ser	UIO	110	Leu	***	VOP					• • •
1381							) TC	CTG	GAA	TCG	GCR	CIG	ACG	GTA	AGC	GAG	cro	GTA	AGC	ACG	1440
1381	GAA	GCC	CTA	AAA	Ala	Lva	Tla	Lau	Glu	Sar	gly	Leu	Thr	Val	Ser	Glu	Leu	Val	Ser	Thr	480
1441	GCA	TGG	cci	TCT	GCA	TCT	ACT	TIT	AGA	AAC	TCT	GAC	AAG	CGC	GGC	OGT	acc	AAC	OCT	GCA	1500
481	Ala	Trp	Ala	Ser	Ala	Ser	Thr	Phe	Arg	Asn	Ser	Asp	Lys	Arg	aly	Gly	Ala	λen	Gly	Alz	500
																					15/0
1501	ccr .	ATA	CGA	CTG	GCC	CCA	CAA	AAA	GAC	TGG	GAA	GTA	AAC	AAC	car	CIAG	CAA	CTT	gcc	AGG	1560 520
501	Arg	Il•	Arg	Leu	Ala	Pro	Gln	Lys	Хвр	Trp	Olu	Val	Asn	Yeu	S.C.O	GIN	OIR	Leu	~~~	vrd	320
											<b>-</b> 1.0		225	cna	aca	CAA	TCA	GAT	AAC	AAA	1620
1561	GTA Val	CIC	AAA	ACA	CTA	GAA	GGI	AIC	Gla	Glu	Ann	Phe	Ann	Gln	Ala	Gln	Ser	λsp	Asn	Lye	540
1621	cm	272	TCG	TTG	GCC	GAC	cro	ATT	GTG	CIG	GCC	<b>GGC</b>	TOT	aca	CCT	OTA	GAA	ጸጸጸ	CCT	GCA	1680
541	Ala	Val	Ser	Lau	Ala	Asp	Leu	Ile	Val	Leu	Ala	Gly	Сув	Ala	GΙΆ	Val	Glu	Lys	Ala	Ala	560
1631	AAA	GAT	GCT	GGC	CAT	GAG	GIG	CAG	GTG	CCT	TIC	AAC	ccc	CGA	CCA	GCG	GAT	GCC	ACC	GCT Bla	1740 530
261	Lys .	Asp	Ala	CJĀ	His	Glu	Val	Gln	Val	Pro	Phe	Ast.	Pro	GIA	Arg	YTA	Yab	W14	1111	VT.	340
													cci	ccc	cc+	CAC	ccc	TIT	AGA	AAC	1800
1741	GAG Glu	CAA	ACC	GAT	GTG	CAA	GCI	TIC	GAA	GCA	CIA	GAG.	DrA	31a	Ala	Ano	Glv	Phe	Arq	Asn.	600
581	Glu	Gln	Thr	Yab	VAL	Gra	WIT	LUG	GIU	~-			•••			•	-				
	TAC .	. ~~		-cc	GAG	CAT	AAA	GTA	TCC	cc1	CAG	GAA	ATG	CIC.	GTA	GAC	CGG	GCG	CAG	CII	1860
1801	T/=	Tia.	Lva	Pro	Glu	His	Lys	Val	Sez	Ala	Glu	Glu	Mac	Leu	Val	qtá	スニタ	Ala	Gln	Leu	620
1461	CIG	TCG	CIT	TCG	GCA	CCX	GAA	ATG	ACT	oc:	TIG	στλ	GGC	OGI	ATO	car	GIA	crc.	GGC	ACC	1920 640
621	Lou	Ser	Leu	Ser	Ala	5L0	Glu	Het	Thr	Y) =	Leu	Val	Gly	Gly	Mec	Arg	ANT	Leu	GIY	1.11	•40
									~~~		3.03		330	cca	CCT	CAG	CTA	TCC	AAT	GAC	1980
1921	AAC Asn	TAC	CYC	GGT	103	CAG	CAT	GGA	G10	111	The	200	Tare.	Pro	Gly	Gln	Leu	Ser	Asn	Asp	660
641	Asn	Tyr	Хвр	GIY	Ser	GIN	HIS	GIY	441	70	****		_, -		•						
	TIC			110		CT3	ga c	crc	AAC	ACT	X AX	TOG	CCIA	acc	AGC	GAT	CAA	TCA	GAC	AAA	2040
1981	Phe	The	Val	AAC	Leu	Leu	Asp	Leu	Ann	Thr	Lys	Trp	Arg	Ala	Ser	Asp	Glu	Ser	yeb	Lys	680
661																					
2041	GII	TTT	CAA	GGC	AGA	CAC	TTC	AAA	ACT	GGC	GAA	GTA	AAG	TOG	agt	occ	ACC	ccc	OIA	CAC	2100
681	Val	Phe	Glu	gly	Yza	λσр	Phe	Lya	Thr	Oly	Gļu	Val	Lys	Trp	Ser	grå	Thr	vid	Val	v=b	700
																					2160
2101	CTO	ATC	TTC	CCA	TCC	AAT	TCC	GAG	CTA	AGA A	Ala	Lau	AL A	Glu	Val	Tyt	01y	Cys	Ala	λ øp	720
701																					
	TCT	Ca A	ON A	3 8/	لململ ا	017	, AAA	CAT	TII	ara	AXG	acc	TGG	acc	ж	GTA	ATG	GAC	CTO	GAC	2220
721	Ser	GI.	G)	Lve	Phe	Val	Lye	λep	2he	Val	Lye	Ala	Trp	λla	Lye	Val	Hot	λsp	Leu	Asp	740
/21	201						•	-													
2221	cca	TTI	GAT	cro	AAA 6	AKT .	. 2	238													
***					Lve			46													

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/16513

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A. CL	ASSIFICATION OF SUBJECT MATTER :C12N 9/08, 15/53, 15/63, 1/21, 15/09; C12P 1/0	0: C12O 1/30									
US CL :435/192, 320.1, 252.3, 41, 27; 536/23.2											
	to International Patent Classification (IPC) or to bo	th national classification and IPC									
	LDS SEARCHED										
	documentation searched (classification system follow	ed by classification symbols)									
U.S. :	435/192, 320.1, 252.3, 41, 27; 536/23.2										
Document	ttion searched other than minimum documentation to the	ne extent that such documents are included	d in the fields searched								
	data base consulted during the international search (ree Extra Sheet.	name of data base and, where practicable	le, search terms used)								
C. DOC	UMENTS CONSIDERED TO BE RELEVANT										
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.								
X	FORKL H. et al. Molecular Cloni Expression of the Gene for Catalase-		3, 13								
A	Photosynthetic Bacterium Rhodobact	` ' '	1, 2, 4-9, 14-17								
	Biochem. 1993, Vol. 214, pages 251-	-									
LOPRASERT, S. et al. Cloning, Nucleotide Sequence, and 3, 13											
	Expression in Escherichia coli of the Bacillus stearothermophilus										
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Furth	er documents are listed in the continuation of Box (C. See patent family annex.									
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		combined with one or more other such being obvious to a person skilled in t	documents, such combination								
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orm PCT/IS	SA/210 (second sheet)(July 1992) *		\ <i>i</i>								

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/16513

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, MEDLINE, SCISEARCH, LIFESCI, EMBASE, WPI, CAS, NTIS, BIOTECHDS, BIOSIS search terms: catalase#, acaligenes or delaya or aquamarinus, microscilla or furvescens

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claims 1-9 and 13-17, drawn to catalases, method of making and method of use thereof. Group II, claims 10-12, drawn to catalase antibodies.

The inventions listed as Groups I and II do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they tack the same or corresponding special technical features for the following reasons: the proteins of Groups I and II are structurally unrelated amino acid sequences.